\* GDVEKGKKIFIMKCSQCHTVEGGKHKTGPNLHGLFGRK SEQ ID NO: 1 human

SEQ ID NO: 2 bacterium: EGDAAAGEKVSKKCLACHTFDQGGANKVGPNPNLFGVF

: GD[x3.3]G[x0.1]K[x0.2]K[x4.0]KC[x2.2]CHT[x3.3]GG[x2.2]K **L**CS

GD{x1.4}E{x3.2}K{x0.2}K{x0.4}KC{x2.2}CHT{x3.3}GG{x2.2}K

homology:47%

Rat : MSLAILRVIRLVRVFRIFKLSRHSKGLQILGRTLKASMRELGLLIFFIGVV

SEQ ID NO: 3

leucinzip, L{6}L{6}L{6}L{6}L

SEQ ID NO: 1 SEQ ID NO: 2 human : GDVEK G K KIFIMKCSQCHTVEKGG KHKTGPNLHGLFGRK ... bacterium: E GDAAAGEKVSK KCLACHTFDQGGANKV GPNPN LFGVF ...

#### Fig. 23 A

```
AFSLFDKDGD
  TEEQIAEFKE
                VMRSLGQNPT
  GTITTKELGT
                VDADGNGTID
                KMKDTDSEEE
  FPEFLTMMAR
                DGNGYISAAE
  IREAFRVFDK
81
                KLTDEEVDEM
101 LRHVMTNLGE
                QVNYEEFVQM
  IREANIDGDG
121
  MTA
```

AMINO ACID SEQUENCE OF CALMODULIN (EXCERPT FROM PDB)

SEQ ID NO: 4

#### Fig. 23 B

1	AMD'QQAEARA FLSEEM	
21	KAAFDMFDAD GGGDIS	<b>LKEL</b>
4.1	GTVMRMLGQN PTKEEL	IIAC
	EEVDEDGSGT IDFEEFL	_ V M
81	VRQMKEDAKG KSEEEL	ADCF
	RIFDKNADGF IDIEEL	GEIL
		KDSD
1	KNNDGRIDFD EFLKMM	EGVQ
161		

AMINO ACID SEQUENCE OF TROPONIN C SEQ ID NO: 5 (EXCERPT FROM PDB)

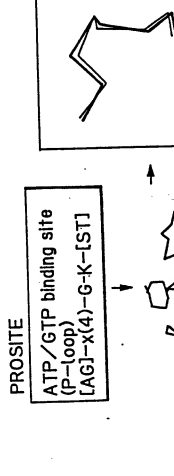
Probe site = 81-108 in Calmodulin

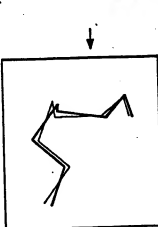
< probe > SEQ ID NO: 6 target probe Sea 10 NO: 7 96 97 98 99 100 101 102 103 104 105 106 107 108 109 110 rmsd = 0.567034

> SEO ID NO: 9 target > SEQ ID NO: 8 < target > SEQ ID NO. 6 > SEQ ID NO: 7 SEQ ID NO: 9 SEQ ID NO: 7 SEQ ID NO: 8 SEQ ID NO: 6 target < target probe probe < probe probe 145 96 97 98 99 100 101 102 103 104 105 106 107 108 109 110 58 144 Probe site = 81-108 and 117-143 in Calmodulin 120 120 150 135

rmsd = 0.823665

```
ensessessesses ATP/GTP binding site exercessessesses
                                                                                                                                                                                                                                                                                                                                          SEQ ID NO: 12
SEQ ID NO: 10
                                                                                         SEQ ID NO: 10
                                                                                                                                                        SEQ ID NO: 11
SEQ ID NO: 10
                                                                                                                                                                                                                                                                                                                        10 11 12 13 14 15 16 17
G A G G V G K S < target >
G H V D H G K T < probe >
rmsd=0.421770 rds protein
                                                                                           < probe >
                                   Probe = (elongation factor)
                                                                                                                                              10 11 12 13 14 15
p G S G K G
V D H G K T
                                                                          7 8.9 10 11 12 13 14
G H V D H G K T
                                                                                                                                                                                                                               unit - A
```





7 8 9 10 11 12 13 14 G H V D H G K T

**SEQ ID NO: 10** 

rmsd=0.42[Å]

ELOGATION FACTOR (LEFM)

rmsd=0.46[A]

SEQ 10 NO: 13 ADENYLATE KINASE(3ADK)

#### Fig. 38 A

```
TVPYQVSLNS
   IVGGYTCCAN
             S
   GYHFC
          G
            G
21
          IQVRL
         G
                    IVHPS
                    KSAAS
         IML
81
                      AGT
              S
            T
101
                    YPD
            SGTS
        T
121
                      YPGQI
   PILSDSSCKS
141
       GYLEGGK
                     D S
161
                    VSWGSGCAQK
       SGKLQGI
181
                     NYV
                         SW
            T
              KVC
        GVY
   N K.P
201
               SEQ ID NO: 14
   ASN
221
```

AMINO ACID SEQUENCE OF TRYPSIN (EXCERPT FROM PDB)

#### Fig. 38 B

```
SWPSQISL
   VVGGTEAQRN
                         LIRQN.WV
        SWAHTC
   RSG
       S
21
            VDRE
             TEQ
61
                      YD
         T
                      G
101
                      WG
        SPC
121
                     PTVD
141
                            GGDG
                    SMVC
   SSYWGST
161
                     PLHC
   RSGCQGDSG
181
                     RLGCNV
           TSFVS
        G V
   AVH
201
                     s w I
        TRVSAYI
221
                SEQ ID NO: 15
```

AMINO ACID SEQUENCE OF ELASTASE (EXCERPT FROM PDB)

### Fig. 39 A

```
Key site number 36 - 41 in Trypsin

41 42 43 44 45 46

M T A A H C \langle trypsin

V S A A H C \langle probe \rangle SEQID NO: 16

V S A A H C \langle probe \rangle SEQID NO: 17

d = 12.070038 [A]

r.m.s.d. = 0.061077 [A]

The number of atoms in a probe = 6

The number of atoms in PDB = 240

The number of combination = 1

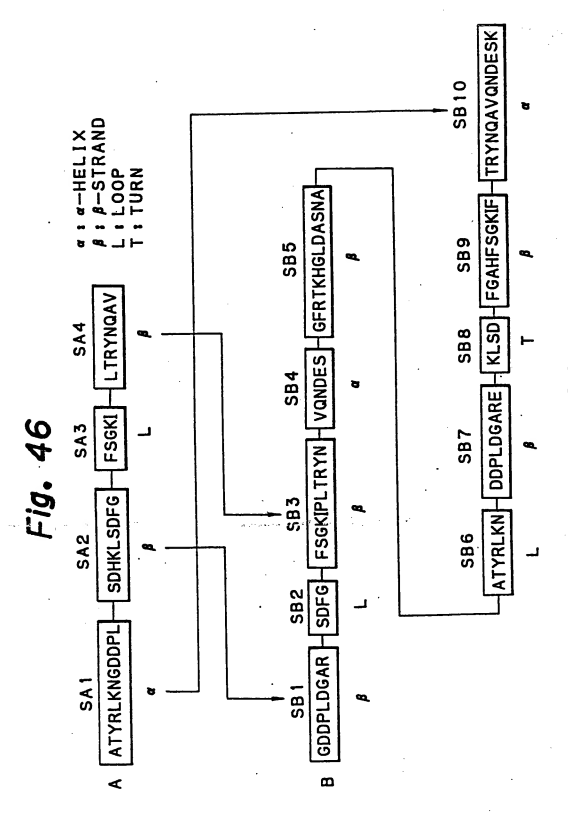
Time = 1sec
```

RETRIEVED RESULTS OF HISTIDINE ACTIVE SITES

### Fig. 39 B

```
Key site number 175 - 179 in Trypsin
186 187 188 189 190
                  G (target > SEQ ID NO: 18
                  G ( probe > SEQ ID NO: 19
              G
          S
      D
  G
              G
          S
      D
  G
d = 8.922721 [A]
r.m.s.d. = 0.092879 [A]
The number of atoms in a probe = 5
The number of atoms in PDB = 240
 The number of combination = 1
 Time = 1sec
```

RETRIEVED RESULTS OF SERINE ACTIVE SITES



SEQ ID NO: 20

SEGIDAN: 1 \* GDVEKGKKIFIMKCSQCHTVEGGKHKTGPNLHGLFGRK human

bacterium : EGDAAAGEKVSKKCLACHTFDQGGANKVGPNPNLFGVF

SEBIDNO: 2

: GD(x3.3}G(x0.1]K(x0.2]K(x4.0]KC(x2.2]CHT(x3.3}GG(x2.2]K LCS

GD{x1,4}E{x0,2}K{x0,2}K{x0,4}KC{x2,2}CHT{x3,3}GG{x2,2}K

homology:47%

: MSLAILRVIRLVRVFRIFKLSRHSKGLQILGRTLKASMRELGLLIFFIGVV Rat

SEQ ID NO: 3

Leucinzip. L{6}L{6}L{6}L{6}L

human : GDVEK G K KIFIMKCSQCHTVEKGG KHKIGPNLHGLFGKN ... SAG I DAG TONDA : E GDAAAGEKVSK KCLACHTFDQGGANKV GPNPN LFGVF... SAG I DAG GDVEK G K KIFIMKCSQCHTVEKGG KHKTGPNLHGLFGRK ... SARTDANS

#### Fig. 23 A

```
AFSLFDKDGD
  TEEQIAEFKE
  GTITTKELGT
               VMRSLGQNPT
  EAELQDMINE
               VDADGNGTID
  FPEFLTMMAR
               KMKDTDSEEE
               DGNGYISAAE
  IREAFRVFDK
101 LRHVMTNLGE
               KLTDEEVDEM
               QVNYEEFVQM
  IREANIDGDG
  MTA
141
```

AMINO ACID SEQUENCE OF CALMODULIN SEQ ID NO: 4 (EXCERPT FROM PDB)

#### Fig. 23 B

1	AMD'QQAEARA	FLSEEMIAEF
21	KAAFDMFDAD	GGGDISTKEL
4.1	GTVMRMLGQN	PTKEELDAII
61	EEVDEDGSGT	IDFEEFLVM
81	VRQMKEDAKG	KSEEELADCF
101	RIFDKNADGF	IDIEELGEIL
121	RATGEHVTEE	DIEDLMKDSD
141	KNNDGRIDFD	EFLKMMEGVQ
161	· .	

AMINO ACID SEQUENCE OF TROPONIN C SEQUENCE OF TROPONIN C

\$\$@ \$\$ λθ: 6 < target > 560 x 8 20: 7 < probe > 550 50 40:6 target > probe 96 97 98 99 100 101 102 103 104 105 106 107 108 109 110 120 Probe site = 81-108 in Calmodulin rmsd = 0.567034

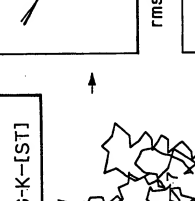
Probe site = 81-108 and 117-143 in Calmodulin

target > scarban < target > sources >55050000 > \$50.Eb scarbas:8 < probe seq 200: 6 SEQ TD 100: 6 < target < probe
556 x b Mb: 3</pre> probe 011 601 801 201 901 144 157 |42 155 4 104 105 54 40 139 153 96 97 98 99 100 101 102 103 137 150 36 135 = 0.823665rmsd

```
55Q ED 40:10
                                                                                                                                                                                                                                                                                                         21:0N RX B555
                                                                                                                                                                   5£9 ED 401 10
                                                                                                                                         SEB ID NO: 11
                                                                                 10 11 12 13 14 15 16 17
G A G G V G K S < target >
G H V D H G K T < probe >
rmsd=0.421770 ras protein
                                                                                                                                8 9 10 11 12 13 14 15
G A P G S G K G < target >
G H V D H G K T < probe >
rmsd=0.648732 adenylate kinase
                            Probe = (elongation factor)
                                                                 7 8.9 10 11 12 13 14
G H V D H G K T
```

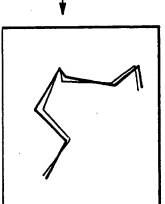
### PROSITE

ATP/GTP binding site (P-loop)
[AG]-x(4)-G-K-[ST]

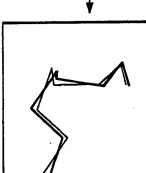


7 8 9 10 11 12 13 14 G H V D H G K T ELOGATION FACTOR (LEFM)

Serbau: 10



rmsd=0.42[Å]



rmsd=0.46[Å]



10 11 12 13 14 15 16 17 G A G G V G K S RAS PROTEIN (2p21)

559 xb no: 12

29/45



15 16 17 18 19 20 21 22 G G P G S G K G 550x3209:13

ADENYLATE KINASE(3ADK)

#### Fig. 38 A

```
IVGGYTCCAN
                  TVPYQVSLNS
                  NSQWVVSAAH
   GYHFCGGSLI
21
   CYKSGIQVRL
                  GEDN
41
   NEQFISASKS
                  IVHPSY
                  KSAASL
   ASISLPTSCA
                  SAGTQC
101
          SGTS
                  YPDVLK
   WGNTKS
121
       SDSSCKS
                  AYPGQI
141
   FCAGYLEGGK
                  DSCQGD
161
   V V C S G K L Q G I
                  VSWGSGCAQK
181
   NKPGVYTKVC
                  NYVSWIKQTI
201
          SEQID NO:14
221
   ASN
```

AMINO ACID SEQUENCE OF TRYPSIN (EXCERPT FROM PDB)

#### Fig. 38 B

```
SWPSQISLQY
  VVGGTEAQRN
                 GGTLIRQNWV
  RSGSSWAHTC
21
                 LTFRVVVGEH
  MTAAHCVDRE
41
                   GVQK
  NLNQNNGTEQ
61
                 GYDIAL
  PYWNTDDVAA
81
  QSVTLNSYVQ
101
                 GWGLTRTN
  LANSPCYITT
121
                 PTV
141
                SMVCAGGDGV
  SSYWGSTVKN
161
  RSGCQGDSGG
                 PLH
181
                 RLGCNVTRKP
   AVHGVTSFVS
201
                 SWINNVIASN
   TVFTRVSAYI
221
```

e com at a s

AMINO ACID SEQUENCE OF ELASTASE (EXCERPT FROM PDB)

#### Fig. 39 A

```
Key site number 36-41 in Trypsin

41 42 43 44 45 46

M T A A H C \langle target \rangle > 560 \text{ Tb AB: 16}

V S A A H C \langle probe \rangle > 560 \text{ Tb AB: 17}

d = 12.070038 [A]

r.m.s.d. = 0.061077 [A]

The number of atoms in a probe = 6

The number of atoms in PDB = 240

The number of combination = 1

Time = 1sec
```

RETRIEVED RESULTS OF HISTIDINE ACTIVE SITES

#### Fig. 39 B

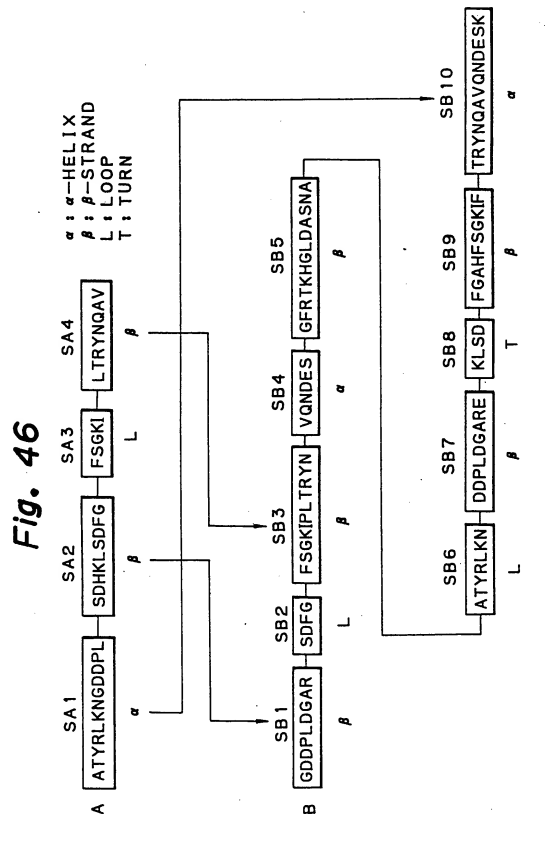
```
Key site number 175 — 179 in Trypsin

186 187 188 189 190

G D S G G \langet \rangle \subseteq \tau \rangle \r
```

RETRIEVED RESULTS OF SERINE ACTIVE SITES

Ĺ



550 TD NO: 20